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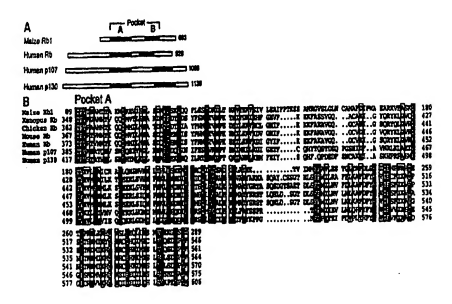
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(54) Title: PLANT RETINOBLASTOMA-ASSOCIATED PROTEINS



#### (57) Abstract

The present invention is based on the isolation and characterization of a plant cell DNA sequence encoding for a retinoblastoma protein. Such finding is based on the structural and functional properties of the plant retinoblastoma protein as possible regulator of the cellular cycle, of the cellular growth and of the plant cellular differentiation. For this reason, among other aspects, it is claimed the use of retinoblastoma protein or the DNA sequence which encodes for it in the growing control of vegetable cells, plants and/or vegetable virus, as well as the use of vectors, cells, plants or animals, or animal cells modified through the manipulation of the control route based on plant retinoblastoma protein.

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# PLANT RETINOBLASTOMA-ASSOCIATED PROTEINS DESCRIPTION

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The present invention relates the proteins having biological activity in plant and animal systems, to polynucleotides encoding for the expression of such proteins, to oligonucleotides for use in identifying and synthesizing these proteins and polynucleotides, to vectors and cells containing the polynucleotides in recombinant form and to plants and animals comprising these, and to the use of the proteins and polynucleotides and fragments thereof in the control of plant growth and plant vulnerability to viruses.

Cell cycle progression is regulated by positive and negative effectors. Among the latter, the product of the retinoblastoma susceptibility gene (Rb) controls the passage of mammalian cells through G1 phase. In mammalian cells, Rb regulates G1/S transit by inhibiting the function of the E2F family of transcription factors, known to interact with sequences in the promoter region of genes required for cellular DNA replication (see eg Weinberg, R.A. Cell 81,323 (1995); Nevins, J.R. Science 258,424 (1992)). DNA tumor viruses that infect animal cells express oncoproteins that interact with the Rb protein via a LXCXE motif, disrupting Rb-E2F complexes and driving cells into S-phase (Weinberg ibid; Ludlow, J. W. FASEB J. 7, 866 (1993); Moran, E. FASEB J. 7, 880 (1993); Vousden, K. FASEB J. 7, 872 (1993)).

The present inventors have shown that efficient replication of a plant geminivirus requires the integrity of an LXCXE amino acid motif in the viral RepA protein and that RepA can interact with members of the human Rb family in yeast (Xie, Q., Suárez-López, P. and Gutiérrez, C. EMBO J. 14, 4073 (1995). The presence of the LXCXE motif in plant D-type cyclins has also been reported (Soni, R., Carmichael, J. P., Shah, Z. H. and Murray, J.

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A. H. Plant Cell 7, 85-103 (1995)).

identified inventors have now The present characteristic sequences of plant Rb proteins corresponding encoding polynucleotides for the first time, isolated such a protein and polynucleotide, and particularly have identified sequences that distinguish it from known animal Rb protein sequences. The inventors have determined that a known DNA sequence from the maize encoding a vegetable Rb plant protein and is hereinafter called ZmRb1. ZmRb1 has been demonstrated by inventors to interact in yeasts with RepA, a plant qeminivirus protein containing LXCXE motif essential for its function. The inventors have further determined that geminivirus DNA replication is reduced in plant cells transfected with plasmids encoding either ZmRb1 or human p130, a member of the human Rb family.

Significantly the inventors work suggests that plant and animal cells may share fundamentally similar strategies for growth control, and thus human as well as plant Rb protein such as ZmRb1 will be expected to have utility in, inter alia, plant therapeutics, diagnostics, growth control or investigations and many such plant proteins will have similar utility in animals.

In a first aspect of the present invention there is provided the use of retinoblastoma protein in controlling the growth of plant cells and/or plant viruses. Particularly, the present invention provides control of viral infection and/or growth in plant cells wherein the virus requires the integrity of an LXCXE amino acid motif in one of its proteins, particularly, e. g., in the viral RepA protein, for normal reproduction. Particular plant viruses so controlled are Geminiviruses.

A preferred method of control using such proteins involves applying these to the plant cell, either directly or by introduction of DNA or RNA encoding for

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their expression into the plant cell which it is desired to treat. By over expressing the retinoblastoma protein, or expressing an Rb protein or peptide fragment thereof that interacts with the LXCXE motif of the virus but does not affect the normal functioning of the cell, it is possible to inhibit normal virus growth and thus also to produce infection spreading from that cell to its neighbours.

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Alternatively, by means of introducing anti-sense DNA or RNA in plant cells in vectors form that contain the necessary promoters for the DNA or RNA transcription, it will be possible to exploit the well known anti-sense mechanism in order to inhibit the expression of the Rb protein, and thus the S-phase. Such plants will be of use, among other aspects to replicate DNA or RNA until high levels, e.g. in yeasts. The methods to introduce anti-sense DNA in cells are very well known for those skilled in the art: see for example "Principles of gene manipulation - An introduction to Genetic Engineering (1994) R.W. Old & S.B. Primrose; Oxford-Blackwell Scientific Publications Fifth Edition p398.

In a second aspect of the present invention there is provided recombinant nucleic acid, particularly in the form of DNA or cRNA (mRNA), encoding for expression of Rb protein that is characteristic of plants. This nucleic acid is characterised by one or more characteristic regions that differ from known animal Rb protein nucleic acid and is exemplified herein by SEQ ID No 1, bases 31-2079.

The DNA or RNA can have a sequence that contains the degenerated substitution in the nucleotides of the codons in SEQ ID No. 1, and in where the RNA the T is U. The most preferred DNA or RNA are capable of hybridate with the polynucleotide of the SEQ ID No. 1 in conditions of low stringency, preferably being the hybridization

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produced in conditions of high stringency.

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The expressions "conditions of low stringency" and "conditions of high stringency" are understood by those skilled, but are conveniently exemplified in US 5202257, Col-9-Col 10. If some modifications were made to lead to the expression of a protein with different amino acids, preferably of the same kind of the corresponding amino acids to the SEQ ID No 1; that is, are conservative substitutions. Such substitutions are known by those skilled, for example, see US 5380712, and it is only contemplated when the protein has activity with retinoblastoma protein.

Preferred DNA or cRNA encodes for a plant Rb protein having A and B pocket sub-domains having between 30% and 75% homology with human Rb protein, particularly as compared with p130, more preferably from 50% to 64% homology. Particularly the plant Rb protein so encoded has the C706 amino acid of human Rb conserved. Preferably the spacer sequence between the A and B pockets is not conserved with respect to animal Rb proteins, preferably being less than 50% homologous to the same region as found in such animal proteins. Most preferably the protein so encoded has 80% or more homology with that of SEQ NO 2 of the sequence listing attached hereto, still more preferably 90% or more and most preferably 95% or more. Particularly provided is recombinant DNA of SEO ID No 1 bases 31 to 2079, or the entire SEQ ID No 1, or corresponding RNAs, encoding for maize cDNA clone encoding ZmRb1 of SQ ID No 2.

In a third aspect of the present invention there is provided the protein expressed by the recombinant DNA or RNA of the second aspect, novel proteins derived from such DNA or RNA, and protein derived from naturally occurring DNA or RNA by mutagenic means such as use of mutagenic PCR primers.

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In a fourth aspect there are provided vectors, cells and plants and animals comprising the recombinant DNA or RNA of correct sense or anti-sense, of the invention.

In a particularly preferred use of the first aspect there is provided a method of controlling cell or viral growth comprising administering the DNA, RNA or protein of the second or third aspects to the cell. Such administration may be direct in the case of proteins or may involve indirect means, such as electroporation of plant seed cells with DNA or by transformation of cells with expression vectors capable of expressing or over expressing the proteins of the invention or fragments thereof that are capable of inhibiting cell or viral growth.

Alternatively, the method uses an expression vector capable of producing anti-sense RNA of the cDNA of the invention.

Another one of the specific characteristics of the plants protein and of the nucleic acids includes a N-terminal domain corresponding in sequence to the amino acids 1 to 90 of the SEQ ID No. 2 and a nucleotides sequence corresponding to the basis 31 to 300 of the SEQ ID No. 1. These sequences are characterized by possessing less than 150 and less than 450 units that the animal sequences which possess more than 300 amino acids and 900 pairs of more bases.

The present invention will now be illustrated further by reference to the following non-limiting Examples. Further embodiments falling within the scope of the claims attached hereto will occur to those skilled in the light of these.

Figures.

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Fig. 1. The sub-figure A shows the relative lengths of the present ZmRb1 protein and the human retinoblastoma proteins. The sub-figure B shows the alignment of the amino acids sequences of the Pocket A and Pocket B of the ZmRb1 with that of the Xenopus, chicken, rat and three human protein (Rb, p107 and p130).

Fig. 2. This figure is a map of the main characteristics of the WDV virus and the pWori vector derived from WDV and the positions of the deletions and mutations used in order to establish that the LXCXE motif is required for its replication in plants cells.

#### EXAMPLE 1.

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10 <u>Isolation of DNA and protein expressing clones.</u>

Total RNA was isolated from maize root and mature leaves by grinding the material previously frozen in liquid nitrogen essentially as described in Soni et al (1995). The major and minor p75ZmRb1 mRNAs were identified by hybridization to a random-primed 32P-labelled PstI internal fragment (1.4 kb).

A portion of a maize cDNA library (106 pfu) in 1ZAPII (Stratagene) was screened by subsequent hybridization to 5'-labelled oligonucleotides designed to be complementary to a known EST sequence of homologue maize of p130. These oligonucleotides were 5'-AATAGACACATCGATCAA/G (M.5m, nt positions 1411-1438) and 5'-GTAATGATACCAACATGG (M.3c, nt positions 1606-1590) (Isogen Biosciences).

After the second round of screening, pBluescript SK-(pBS) phagemids from positive clones were isolated by in vivo excision with ExAssist helper phage (Stratagene) according to protocols recommended by the manufacturer. DNA sequencing was carried out using a SequenaseTM Kit (USB).

The 5'-end of the mRNAs encoding p75ZmRbl was determined by RACE-PCR. Poly-A+mRNA was purified by chromatography on oligo-dT-cellulose (Amersham). The first strand was synthesized using oligonucleotide DraI35 (5'-GATTTAAAATCAAGCTCC, nt positions 113-96). After denaturation at 90°C for 3 min, RNA was eliminated by

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RNase treatment, the cDNA recovered and 5'-tailed with terminal transferase and dATP. Then a PCR fragment was amplified using primer DraI35 and the linker-primer (50 bp) of the Stratagene cDNA synthesis kit.

One of the positive clones so produced contained a ~4 kb insert that, according to restriction analysis, extended both 5' and 3' of the region contained in the Expressed Sequence Tag used. The nucleotide sequence corresponding to the longest cDNA insert (3747 bp) is shown in SEQ ID No. 1. This ZmRb1 cDNA contains a single open reading frame capable of encoding a protein of 683 amino acids (predicted Mr 75247, p75ZmRb1) followed by a 1646 bp 3'-untranslated region. Untranslated regions of similar length have been also found in mammalian Rb cDNAs (Lee, W.-L. et al, Science 235, 1394 (1987); Bernards, R. et al, Proc. Natl. Acad. Sci. USA 86, 6474 (1989)). Northern analysis indicates that maize cells derived from both root meristems and mature leaves contain a major message, ~2.7±0.2 kb in length. In addition, a minor Heterogeneous appears. message also  $~3.7\pm0.2$ kb transcripts have been detected in other species (Destrée, O. H. J. et al, Dev. Biol. 153, 141 (1992)).

Plasmid pWoriAA was constructed by deleting in pWorimost of the sequences encoding WDV proteins (Sanz and Gutierrez, unpublished). Plasmid p35S.Rb1 was constructed by inserting the CaMV 35S promoter (obtained from pWDV3:35SGUS) upstream of the ZmRb1 cDNA in the pBS vector. Plasmid p35S.130 was constructed by introducing the complete coding sequence of human p130 instead of ZmRb1 sequences into p35S.Rb1. Plasmid p35.A+B was constructed by substituting sequences encoding the WDV RepA and RepB ORFs instead of ZmRb1 in p35S.Rb1 plasmid. (See Soni, R. and Murray, J. A. H. Anal. Biochem. 218, 474-476 (1994)).

The sequence around the methionine codon at nucleotide

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in maize p75ZmRb1.

position 31 contains a consensus translation start (Kozak, M. J. Mol. Biol. 196, 947 (1987)). To determine whether the cDNA contained the full-length ZmRb1 coding region, the 5'-end of the mRNAs was amplified by RACE-PCR using an oligonucleotide derived from a region close to the putative initiator AUG, which would produce a fragment of ~150 bp. The results are consistent with the ZmRb1 cDNA clone containing the complete coding region.

The ZmRb1 protein contains segments homologous to the A and B subdomains of the "pocket" that is present in all members of the Rb family. These subdomains are separated by a non-conserved spacer. ZmRb1 also contains nonconserved N-terminal and C-terminal domains. Overall, shares ZmRb1 ~28-30% amino acid identity similarity) with the Rb family members (Hannon, G. J., Demetrick, D. & Beach, D. Genes Dev. 7, 2378 (1993); Cobrinik, D., Whyte, P., Peeper, D.S., Jacks, T. & Weinberg, R. A. ibid., p. 2392 (1993). Ewen, M. E., Xing, Y. Lawrence, J. B. and Livingston, D. M. Cell 66, 1155 (1991))(Lee W. L. et al, Science 235, 1394 (1987); Bernards et al, Proc. Natl. Acad. Sci. USA 86, 6974 (1989)), with the A and B subdomains exhibiting the highest homology (~50-64%). Interestingly, amino acid C706 in human Rb, critical for its function (Kaye, F. J., Kratzke R. A., Gerster, J. L. and Horowitz, J. M. Proc. Natl. Acad. Sci. USA 87, 6922 (1990)), is also conserved

Note: The 561-577 amino acids encompass a proline-rich domain.

ZmRb1 contains 16 consensus sites, SP or TP for phosphorilation by cyclins dependant kinases (CDKs) with one of the 5'-tail of the sub-domain A and several in the C-terminal area which are potential sites of phosphorilation. A nucleic acid preferred group which encodes proteins in which one or more of these sites are

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changed or deleted, making the protein more resistant to the phosphorilation and thus, to its functionality, for example linking to E2F or similar. This can be easily carried out by means of mutagenesis conducted by means of PCR.

### EXAMPLE 2

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### In vivo activity.

Replication of wheat dwarf geminivirus (WDV) dependent upon an intact LXCXE motif of the viral RepA protein. This motif can mediate interaction with a member 10 of the human Rb family, p130, in yeasts. Therefore, the inventors investigated whether p75ZmRb1 could complex with WDV RepA by using the yeast two-hybrid system (Fields, S. and Song, O. Nature 340, 245-246 (1989)). Yeast cells were co-transformed with a plasmid encoding 15 the fusion GAL4BD-RepA protein and with plasmids encoding different GAL4AD fusion protein. The GAL4AD-p75ZmRb1 fusion could also complex with GAL4BD-RepA to allow growth of the recipient yeast cells in the absence of histidine. This interaction was slightly stronger than 20 that seen with the human p130 protein. RepA could also bind to some extent to a N-terminally truncated form of p75ZmRb1. The role of the LXCXE motif in RepA-p75ZmRb1 interaction was assessed using a point mutation in WDV RepA (E198K) which we previously showed to destroy 25 interaction with human pl30. Co-transformation of ZmRbl with a plasmid encoding the fusion GAL4BD-RepA(E198K) indicated that the interaction between RepA and p75ZmRb1 occurred through the LXCXE motif.

In this respect, the E198K mutant of WDV RepA behaves similarly to analogous point mutants of animal virus oncoproteins (Moran, E., Zerler, B., Harrison, T. M. and Mathews, M.B. Mol. Cell Biol. 6, 3470 (1986); Cherington, V. et al., ibid., p. 1380 (1988); Lillie, J. W., Lowenstein, P. M., Green, M. R. and Green, M. Cell 50,

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1091 (1987); DeCarpio, J. A. et al., ibid., p. 275 (1988)).

Specific interaction between maize p75ZmRb1 and WDV RepA in the yeast two-hybrid system (Fields et al) relied on the ability to reconstitute a functional GAL4 activity from two separated GAL4 fusion proteins containing the DNA binding domain (GAL4BD) and the activation domain (GAL4AD). Yeast HF7c cells were co-transformed with a plasmid expressing the GAL4BD-RepA or the GAL4BD-RepA(E198K) fusions and the plasmids expressing the GAL4AD alone (Vec) or fused to human p130, maize p75 (p75ZmRb1) or a 69 amino acids N-terminal deletion of p75 (p75ZmRb1-DN). Cells were streaked on plates with or without histidine according to the distribution shown in the upper left corner. The ability to grow in the absence of histidine depends on the functional reconstitution of a GAL4 activity upon interaction of the fusion proteins, since this triggers expression of the HIS3 gene which is under the control of a GAL4 responsive element. The growth characteristics of these yeast co-transformants correlate with the levels of b-galactosidase activity.

Procedures for two-hybrid analysis are described in Xie et al (1995). The GAL4AD-ZmRb1 fusions were construed in the pGAD424 vector.

#### 25 EXAMPLE 3

#### In vivo activity.

Geminivirus DNA replication requires the cellular DNA replication machinery as well as other S-phase specific factors (Davies, J. W. and Stanley, J. Trends Genet. 5, 77 (1989); Lazarowitz, S. Crit. Rev. Plant Sci. 11, 327 (1992)). Consistent with this requirement, geminivirus infection appears to drive non-proliferating cells into S-phase, as indicated by the accumulation of the proliferating cell nuclear antigen (PCNA), a protein which is not normally present in the nuclei of

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differentiated cells (Nagar, S., Pedersen, T. J., Carrick, K. M., Hanley-Bowdoin, L. and Robertson, D. Plant Cell 7, 705 (1995)). The inventors finding that efficient WDV DNA replication requires an intact LXCXE motif in RepA coupled with the discovery of a plant homolog of Rb supports the model that, as in animal cells, sequestration of plant Rb by viral RepA protein promotes inappropriate entry of infected cells into Sphase. Therefore, one way to investigate the function of p75ZmRbl was to measure geminivirus DNA replication in cells transfected with a plasmid bearing the ZmRb1 sequences under a promoter functional in plant cells, an approach analogous to that previously used in human cells (Uzvolgi, E. et al., Cell Growth Diff 2, 297 (1991)). Accumulation of newly replicated viral plasmid DNA was impaired in wheat cells transfected with plasmids expressing p75ZmRb1 or human p130, when expression of WDV replication protein(s) is directed wither by the WDV promoter or by the CaMV 35S promoter.

Since WDV DNA replication requires an S-phase cellular environment, interference with viral DNA replication by p75ZmRb1 and human p130 strongly evidences a role for retinoblastoma protein in the control of the G1/S transition in plants. The existence of a plant Rb homolog implies that despite their ancient divergence, plant and animal cells use, at least in part, similar regulatory proteins and pathways for cell cycle control.

Two lines of evidences reinforce this model. First, a gene encoding a protein that complements specifically the G1/S, but not the G2/M transition of the budding yeast cdc28 mutant has been identified in alfalfa cells (Hirt, H., Páy, A., Bögre, L., Meskiene, I. and Heberle-Bors, E. Plant J. 4, 61 (1993)). Second, plant homologs of D-type cyclins have been isolated from Arabidopsis and these, like their mammalian relatives, contain LXCXE motifs. In

concert with plant versions of CDK4 and CDK6, plant D-type cyclins may regulate passage through G1 phase by controlling the phosphorylation state of Rb-like proteins.

In animal cells, the Rb family has been implicated in 5 tumor suppression and in the control of differentiation and development. Thus, p75ZmRb1 could also play key regulatory roles at other levels during the plant cell life. One key question that is raised by the existence of Rb homologs in plant cells in whether, as in animals 10 disruption of the Rb pathway leads to a tumor-prone condition. In this regard, the inventors have noted that the VirB4 protein encoded by the Ti plasmids of both Agrobacterium tumefaciens and A. rhyzogenes contains an LXCXE motif. Although the VirB4 protein is required for 15 tumor induction (Hooykas, P. J. J. and Beijersbergen, A. M. Annu. Rev. Phytopathol. 32, 157 (1994), the function of its LXCXE motif in this context remains to be examined. Geminivirus infection is not accompanied by tumor development in the infected plant, but in some 20 cases an abnormal growth of enactions has been observed (G. Dafalla and B. Gronenborn, personal communication). Inhibition of wheat dwarf geminivirus replication by ZmRb1 or human p130 in cultured wheat cells was carried out as follows. A. Wheat cells were 25 transfected, as indicated, with pWori (Xie et al. 1995). alone (0.5g), a replicating WDV-based plasmid which encodes WDV proteins required for viral DNA replication, and with control plasmid pBS (10 g) or p35S.Rb1 (10 g), which encodes ZmRb1 sequences under the control of the 30 CaMV 35S promoter. Total DNA was purified one and two days after transfection, equal amounts fractionated in agarose gels and ethidium bromide staining and viral pWori DNA identified by Southern hybridization. Plasmid DNA represents exclusively newly-replicated plasmid DNA 35

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since it is fully resistant to DpnI digestion and sensitive to Mbol. Note that the MboI-digested samples were run for about half of the length than the undigested samples. B. To test the effect of human p130 on WDV DNA replication, wheat cells were co-transfected with pWori (0.5 g) and plasmids pBS (control), p35S.Rb1 or p35S.130 (10 q in each case). Replication of the test plasmid (pWori) was analyzed two days after transfection and was detected as described in part A using ethidium bromide staining; and Southern hybridization. C. To test the effect of ZmRb1 or human p130 on WDV DNA replication when expression of viral proteins was directed by the CaMV 35S promoter, the test plasmid pWoriAA (which does not encode functional WDV replication proteins but replicates when they are provided by a different plasmid, i. e. pWori) was used. Wheat cells were co-transfected, as indicated, with pWori $\Delta\Delta$  (0.25 g), pWori (0.25 g), p35S.A+B (6 g), p35S.Rb1 (10 g) and/or p35S.130 (10 g). Replication of the test plasmid (pWori $\Delta\Delta$ ) was analyzed 36 hours after transfection and was detected as described in part A using ethidium bromide staining; Southern hybridization. Plasmids pWori (M1) and pWoriΔΔ (M2; Sanz and Gutiérrez, unpublished), 100 pg in each case, were used as markers. Suspension cultures of wheat cells, transfection by of viral DNA bombardment and analysis particle replication were carried out as described in (Xie et al. 1995), except that DNA extraction was modified as in (Soni and Murray. Arnal. Biochem. 218, 474-476 (1995).

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### SEQUENCE LISTING

- (1) GENERAL INFORMATION:
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    - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 3747 base pairs
- 25 (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: unknown
  - (ii) MOLECULE TYPE: DNA (genomic)
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- 30 (iv) ANTI-SENSE: NO
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### (B) LOCATION: 31..2079

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

GAAT	TCG(	GCA (	CGAGO	CAAAC	G TO	CTGAT	TGAT					_			T TTG	54
					•			1	l .			9	5			
							<b>-</b>									
				AAA												102
GIu	-	Met	GIu	Lys	Leu	-	Asn	ser	Asn	ser	-	Lys	GIY	GIU	Leu	
	10					15					20					
GAT	TTT	AAA	TCA	ATT	TTG	ATC	AAT	аат	GAT	тат	ATT	CCC	TAT	GAT	GAG	150
Asp	Phe	Lys	Ser	Ile	Leu	Ile	Asn	Asn	Asp	Тут	Ile	Pro	Tyr	Agp	Glu	
25		-			30					35					40	
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naA	Ser	Thr	Gly	qaA	Ser	Thr	naA	Leu	Gly	aiH	Ser	Lyo	Cyp	Ala	Phe	
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				TCT												246
GIU	inr	Leu		Ser	Pro	Inr	гув	65	116	Lys	ASII	Mec	70	Inr	val	
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		75					80					85				
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Gln	Met	Thr	Pro	Val	Thr	Ser	Ala	Met	Thr	Thr	Ala	Lys	Trp	Leu	Arg	
	90					95					100					
									~							
				TCA												390
	vai	ile	ser	Ser	110	Pro	Авр	гÀв	Pro	115	ser	Lys	Leu	GIR	120	
105					110					113					120	
TTT	CTG	TCA	TCA	TGC	GAT	AGG	GAT	TTG	ACA	ААТ	GCT	GTC	ACA	GAA	AGG	438
Phe	Leu	Ser	Ser	Сув	Asp	Arg	Авр	Leu	Thr	Asn	Ala	Val	Thr	Glu	Arg	
				125					130					135		
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Val	Ser	He	Val	Leu	Glu	Ala	Ile	Phe	Pro	Thr	Lys	Sex	Ser	Ala	Asn	
			140				•	145					150			
000	CCT	OTT	TCC	دبيملا	CCC	CTC	חאא	ጥርጥ	CCA	አአጥ	acc	umanı	ara.	y down	ccc	E 3.4
				TTA												534
wr.d	GIY	155	943 I.	Lieu	GIY	·	160	CAR	n1d	U911	n.a	165	valy	116	210	
							- 55									
TGG	GCV	Gλλ	GCC	AGA	AAA	GTG	GAG	GCT	TCC	AAG	TTG	TAC	TAT	AGG	GTA	582
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	170					175					180					

												AAT				630
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												TTT Phe 245				774
												CTC Leu				822
												TTG Leu				870
												ATT 11e				918
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												AAG Lys				1158
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- 17 -

Phe	Ala	Ser 395	Pro	Thr	Val	Cys	Asn 400	Pro	Val	Gly	Gly	Asn 405	Glu	Lyc	Сув	
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Thr	Glu	Arg	Val	Tyr	Asn	Val	Phe	Lys	Gln	Ile	Leu	G1u	Gln	Gln	Thr	
				445					450					455		
ACA	TTA	TTT	TTT	AAT	AGA	CAC	ATC	GAT	CAA	CTT	ATC	CTT	TGC	TGT	CTT	1446
Thr	Leu	Phe	Pho	Asn	Arg	His	lle	Авр	Gln	Leu	Ile	Leu	Cys	Cys	Leu	
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TAT	GGT	GTT	GCA	AAG	GTT	TGT	CAA	TTA	GAA	CTC	ACA	TTC	AGG	GAG	ATA	1494
Tyr	${\tt Gl}_{Y}$	Val	Ala	Lys	Val	Cys	Gln	Leu	Glu	Leu	Thr	Phe	Arg	Glu	Ile	
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Sei	He	Tyr	He	Gly	Ser	Thr	Asn	Arg	naA		Val	Leu	Val	Ser		
505					510					515					520	
						TTT										1638
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ТЛЛ		TAT	CTG	тст	ccr	TTG	CGG	CAA	ACC	AAG	TTG	GAT	CTA	CTG	CT'G	1830
	GTA															1830
	GTA Val					TTG										1830

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Thr Ser Aop Pro Ala Ala Ala I 665 - 670	Phe Ser Pro Leo 675		
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STCTCATGTT GTGACTTCAT TGCTGAT			•
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ITCCCAAATT GCAATTANTC CAGANGT			
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SACATOGON' CTOTTTOTOC AAGTGAG	ATG AAGAAAACCI	GAAATGCTAT CACCATTTAA	3019

- 19 -

AACATTGGCT	TCTGGAAGTT	CAGGTGATTA	GCAGGAGACG	TTCTGACATT	GCCATTGACA	3079
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AGAAGGCGAG	AAGTCATTGC	TGCCAGCATT	GTGATCGTCA	GGCACAGAAG	TACTCAAAAG	3499
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TCTATATACA	AGCAGCGGAT	TTTGTTTAGA	GTTAGCTTTT	GAGATGCATC	ATTTCTTTCA	3619
CATCTGATTC	TGTGTGTTGT	AACTCGGAGT ·	CGCGTAGAAG	TTAGAATGCT	AACTGACCTT	3679
AATTTTCACC	GAATAATTTG	CTAGCGTTTT	TCAGTATGAA	ATCCTIGTCT	ЛААААААА	3739
лаааааа						3747

### (2) INFORMATION FOR SEQ ID NO: 2:

### (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 683 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

### (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

Met Glu Cys Phe Gln Ser Asn Leu Glu Lys Met Glu Lys Leu Cys Asn 1 5 10 15

Ser Asn Ser Cys Lys Gly Glu Leu Asp Phe Lys Ser Ile Leu Ile Asn 26 25 30

Asn Asp Tyr Ile Pro Tyr Asp Glu Asn Ser Thr Gly Asp Ser Thr Asn 35 40 45

Leu Gly His Ser Lys Cys Ala Phe Glu Thr Leu Ala Ser Pro Thr Lys
50 55 60

65	Ile	liye	Asn	Met	Leu 70	Thr	Val	Pro	Ser	Ser 75	Pro	Leu	Ser	Pro	Ala 80
Chr.	Gly	Glγ	Ser	Val 85	Lys	Ile	Val	Gln	Met 90	Thr	Pro	Val.	Thr	Ser 95	Ala
1e: t	Thr	Thr	Ala 100	Lys	Trp	Leu	Arg	Glu 105	Val	Ile	Ser	Ser	Leu 11.0	Pro	Asp
Lys	Pro	Ser 115	Ser	Г'n	Leu	Gln	Gln 120	Phe	Leu	Ser	Ser	Сув 125	qaA	Arg	Дяр
bau	Thr 130	Ásn	Ala	Val	Thr	Glu 135	Arg	Val	Ser	lle	Val 140	Leu	Glu	Ala	Ile
145				Ser	150					155					160
				Pho 165					170					175	
٠			180	Tyr				185					190		
		195		Asn			200					205			
	210			Cys		215					220				
225				Val Phe	230					235					240
				245 Leu					250					255	
			260	Leu				265					270		
		275		ile			280					285			
	290			Leu		295					300				
305				Val	310					315					320
				325 Gly					330					335	
ьys	, vi ĉ	WTG	340		.,,	veħ	voh	345		տոլո	.10	ii	350		276

- 21 -

Arg Ser Cys Asn Glu Ser Arg Asn Thr Val Val Glu Arg Asn Leu Gln 355 360 365

Thr Pro Pro Pro Lyc Gln Ser His Met Val Ser Thr Ser Leu Lys Ala 370 375 380

Lys Cys His Pro Leu Gln Ser Thr Phe Ala Ser Pro Thr Val Cys Asn 385 390 395 400

Pro Val Gly Gly Asn Glu Lys Cys Ala Asp Val Thr Ile His Ile Phe 405 410 415

Phe Ser Lye Ile Leu Lye Leu Ala Ala Ile Arg Ile Arg Asn Leu Cyc 420 425 430

Glu Arg Val Gln Cys Val Glu Gln Thr Glu Arg Val Tyr Asn Val Pho 435 440 445

Lye Gln (le Leu Glu Gln Gln Thr Thr Leu Phe Phe Asn Arg His 1]e 450 455 460

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heu Glu Leu Thr Phe Arg Glu IIe Leu Asn Asn Tyr Lys Arg Glu Ala 485 490 495

Gln Cys Lys Pro Glu Val Phe Ser Ser Ile Tyr Ile Gly Ser Thr Asn 500 505 510

Arg Ash Gly Val Lou Val Ser Arg His Val Gly Ile Ile Thr Phe Tyr 515 520 525

Asn Glu Vai Phe Val Pro Ala Ala Lys Pro Phe Leu Val Ser Leu Ile 530 535 540

 Ser Ser Gly Thr His Pro Glu Asp Lys Lys Asn Ala Ser Gly Gln 11c

 545
 550
 555
 560

Pro Gly Ser Pro Lye Pro Ser Pro Phe Pro Asn Leu Pro Asp Met Ser 565 570 575

Pro Lys hys Val Ser Ala Ser His Ash Val Tyr Val Ser Pro Leu Arg 580 585 . 590

Gln Thr Lys Leu Asp Leu Leu Leu Ser Pro Ser Ser Arg Ser Phe Tyr 595 600 605

Al. Cys lle Gly Glu Gly Thr His Ala Tyr Gln Ser Pro Ser Lys Asp

Leu Ala Ala ile Asn Ser Arg Leu Asn Tyr Asn Gly Arg Lys Val Asn 625 630 635 640

- 22 -

Ser Arg Leu Asn Phe Asp Met Val Ser Asp Ser Val Val Ala Gly Ser 645 650 655

Leu Gly Gln Ile Asn Gly Gly Ser Thr Ser Asp Pro Ala Ala Ala Phe 660 665 670

Ser Pro Leu Ser Lys Lys Arg Glu Thr Asp Thr 675 680

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INFORMATION RELATIVE TO THE DEPOSIT OF A MICRO-ORGANISM
The micro-organism to which reference is made in page
6 of the disclosure has been deposited in the following
institution:

- COLECCION ESPAÑOLA DE CULTIVOS TIPO (CECT)
  Departamento de Microbiología
  Facultad de Ciencias Biológicas
  46100 BURJASOT (Valencia)
  Spain
- Deposit identification: pBS.Rb1
  Deposit date: June 12, 1996
  Order No.: 4699

This information appears reflected in the form PCE/RO/134 enclosed to the request.

## INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism relate on page6	ferred to in the description and following
B. IDENTIFICATION OF DEPOSIT pBS. Rb1	Further deposits are identified on an additional sheet
Name of depository institution COLECCION ESPAÑOLA DE CULTIVOS	
Address of depositary institution (including pastal code and country)  Departamento de Microbiología  Facultad de Ciencias Biológicas  46100 BURJASOT (Valencia)  Spain	
Date of deposit 12 June 1996	Accession Number 4699
C. ADDITIONAL INDICATIONS (leave blank if not applicable	e) This information is continued on an additional shoet
D. DESIGNATED STATES FOR WHICH INDICATION	IS ARE MADE (if the indications are not for all designated States)
E. SEPARATE FURNISHING OF INDICATIONS (leave leave indications listed below will be submitted to the International Enternational	
For receiving Office we only	For International Bureau uso only
This shoet was received with the international application	This sheet was received by the International Bureau on:
Authorized officer  C.A.J.A. PASCHE	Authorized afficer
mm PCT/RO/134 (July 1992)	

### CLAIMS

- Use of a retinoblastoma (Rb) protein for the control of the growth and/or replication of plant cells and plant viruses.
  - 2. Use as claimed in claim 1 characterised in that the virus requires the integrity of an LXCXE amino acid motif in one of its proteins for the normal reproduction.
- 3. Use as claimed in claim 1 wherein the virus is a Geminivirus.
- 4. Use in accordance with claim 1 characterised in that the virus binds a retinoblastoma (Rb) protein in order to release a transcription factor.
- 5. A method of controlling the growth and/or replication of a plant cell or a plant virus within that cell, comprising the increase or decrease of the level and/or activity of a retinoblastoma protein in that plant cell.
  - 6. A method as claimed in claim 5 characterised in that the level of protein is increased by direct application.
  - 7. A method as claimed in claim 5 characterised in that the level of protein is increased by introduction of DNA or RNA encoding for its expression into the plant cell which it is desired to treat.

- 8. A method as claimed in claim 5, 6 or 7 wherein the protein is overexpressed.
- 9. A method of controlling the growth and/or replication of a plant cell or a plant virus comprising expressing an

Rb protein, or peptide fragment thereof that interacts with the LXCXE motif of the virus but does not affect the normal functioning of the cell, such as to inhibit cell growth or normal viral growth.

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- 10. Recombinant nucleic acid encoding for expression of an Rb protein that has one or more characteristics of plant Rb protein not shared by animal Rb protein.
- 10 11. Nucleic acid as claimed in claim 10 characterised in that it comprises one or more characteristic regions that differ from known animal Rb protein nucleic acid.
- 12. Recombinant nucleic acid in the form of DNA or cRNA which encodes for a plant Rb protein having A and B pocket subdomains having a sequence with between 30% and 75% homology with human Rb protein.
- 13. Nucleic acid as claimed in claim 12 having a sequence with between 30% and 75% homology with p130 Rb retinoblastoma protein.
  - 14. Nucleic acid as claimed in claim 12 or 13 characterised in that it has from 50% to 64% homology with animal or p130 Rb retinoblastoma protein.
    - 15. Nucleic acid as claimed in any one of claims 12 to 14 encoding for the C706 amino acid of human Rb.
- 16. Nucleic acid as claimed in any one of claims 12 to 15 wherein the spacer sequence between the A and B pockets is not conserved with respect to animal Rb proteins.
- 17. Nucleic acid as claimed in claim 16 wherein the spacer sequence has less than 50% homology to the same

region found in animal retinoblastoma proteins.

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- 18. Nucleic acid as claimed in any one of claims 12 to 17 having 80% or more homology with that of SEQ NO 2.
- 19. Nucleic acid as claimed in claim 18 wherein the homology is 90% or more.
- 20. Recombinant DNA comprising a sequence corresponding to SEQ ID No 1 bases 31 to 2079. 10
  - 21. Recombinant DNA comprising a sequence corresponding to SEQ ID No 1 or corresponding RNA encoding for maize cDNA clone encoding ZmRb1 of SQ ID No 2.
- 22. Protein encoded by the recombinant DNA or RNA as claimed in any one of claims 12 to 21 or novel proteins derived from such DNA or RNA, and protein derived from naturally occurring DNA or RNA altered by mutagenic means. 20
  - 23. Protein as claimed in claim 22 wherein the mutagenic means comprises mutagenesis using mutagenic PCR primers.
- 24. Anti-sense DNA or RNA of a gene encoding for a plant 25 retinoblastoma protein, a gene which possesses the nucleic acid sequence as the one which is claimed in any one of the claims 10 to 21.
- 25. Vectors, cells, plants or animals comprising the DNA 30 or RNA as claimed in any one of claims 12 to 22.
- growth control and/or method to the 26. A proliferation of a vegetable cell or of a plant virus comprising the decrease of plant retinoblastoma protein 35

levels in the cell by incorporation to this cell of antisense DNA or RNA to the retinoblastoma protein.

- 27. cDNA encoding a protein as it is claimed in the claim 5 22.
  - 28. A nucleic acid encoding a protein in which one or more of these sites are altered or deleted, making the protein more resistant to the phosphorilation and thus, to its functionality, for example, linking to E2F or similar.
    - 29. An encoded protein by the nucleic acid which is described in claim 28.

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Human p107

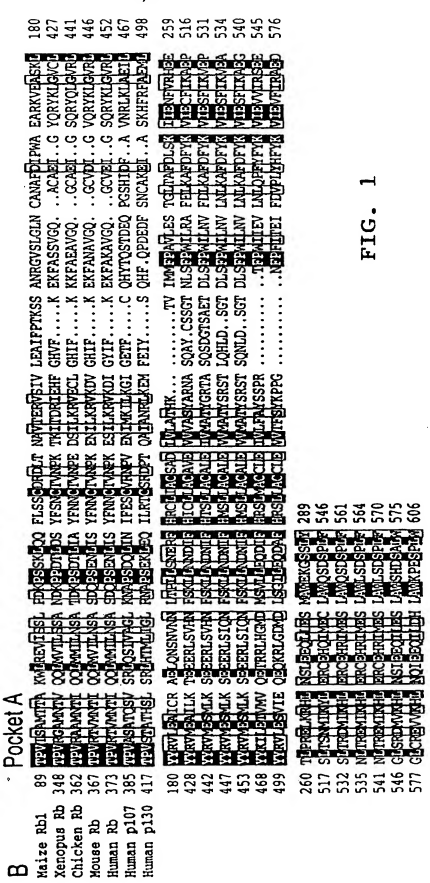
Human Rb

Maize Rb1

Human p130

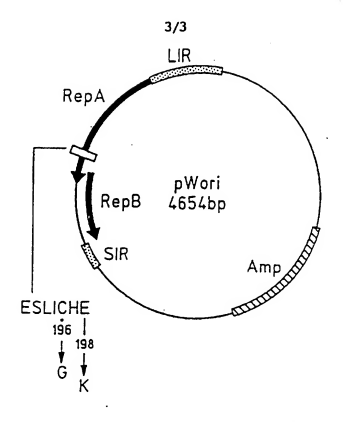
A Pocket B

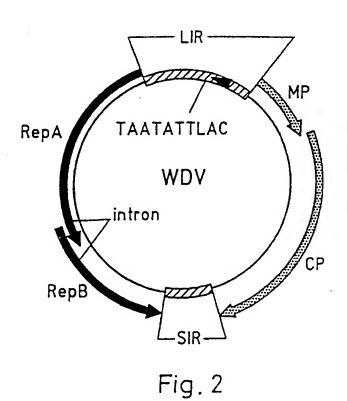




2/3	
492 701 721 728 868 916 916	525 726 746 746 753 931 1006
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	GSTNRNG
HIDOLINGSM HIDOLINGSM HIDOLINGSM HIDOLINGSM HIDOLINGSM HIDOLINGSM HIDOLINGSM HIDOLINGSM	EIG. 1
EQQITIBFFNR QQEYEBMRDR QNEYEBMRDR QNEYEBMRDR VHCPDBMRDR VHCPDBMRDR	PVMRSSSTLP
RYZNVFKQIE VIMILLOHIL LIMILFOHIL IIMILFOHIL IIMILFOHIL KIMICFEFIL	NKDRTSRDSS
VOCVECTE LLSEHPELEO LLSEHPELEH LLSEHPELEH LLSEHPELEH LDVSN.ELRR LDISD.ELRK	REVVAYNKNI RSHQNSPTEL
AIRTRNICER YKRUSSICSS YKRUTUFFR YKRUTUCER SYRUNTICER SYRUNTICER AVRIRDICAK	KRRNSGSSDS
4544444	ESSENTE FKHVLIR FKRVLIR FKRVLIK FKRVLIK FKRVLIK FKRVLIK FKRVLIK FKRVLIK FKRVLIK FRILL 760 LKTNIL 760
Pocket B REKCADVITH COKSTSLS CKPOKSTSLS CKPLKSTSLS CKPLKSTSLS CKPLKSTSLS CKPLKSTSLS CKPLKSTSLS CKPLKSTSLS CKPLKSTSLS CKPLKSTSLS	KREACK PEV KGLTNTNOET KGLTNTNOET KDLPHAVOET KOLPHAVOET RNOPOANSHV BITQPQARSQV TITNEVE MOK VEVNIUVE MOK
405 614 630 632 639 780 828	493 702 720 720 720 917 727 747 747 754 932
Maize Rbi Xenopus Rb Chicken Rb Mouse Rb Human Rb Human p107	

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PCT/EP 97/03070

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 C12N15/29 C12 C12N5/10 C12N15/82 C12N15/11 C07K14/415 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) C07K C12N IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No χ SHEN B. ET AL.: "Partial sequencing and 10-21. mapping of clones from two maize cDNA 25,27 libraries" PLANT MOLECULAR BIOLOGY. vol. 26, no. 4, November 1994, pages 1085-1101, XP002042536 see the whole document "AC T18395" EMBL DATABASE, 23 April 1994, HEIDELBERG, see the whole document Х GRAFI G. ET AL.: "AC U52099" 10-26 EMBL DATABASE. 26 April 1996, HEIDELBERG, XP002042537 Υ see the whole document 1 - 9, 27-/--X Further documents are listed in the continuation of box C. Patent family members are listed in annex. Х \* Special categories of cited documents : \*T\* later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to fding date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or other means nts, such combination being obvious to a person skilled document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 2 3. 10. 97 6 October 1997 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx, 31 651 epo ni, Kania, T Fax: (+31-70) 340-3016

PCT/EP 97/03070

.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
х	QIAN Y ET AL: "BIOLOGICAL FUNCTION OF THE RETINOBLASTOMA PROTEIN REQUIRES DISTINCT DOMAINS FOR HYPERPHOSPHORYLATION AND TRANSCRIPTION FACTOR BINDING" MOLECULAR AND CELLULAR BIOLOGY, vol. 12, no. 12, pages 5363-5372, XP000615356 see the whole document	28,29
×	WO 95 06661 A (RES DEV FOUNDATION ;FUNG YUEN KAI (US)) 9 March 1995 * see especially p.31, first par. *	28,29
Y	XIE Q. ET AL.: "Identification and analysis of a retinoblastoma binding motif in the replication protein of a plant DNA virus: requirement for efficient viral replication"  THE EMBO JOURNAL, vol. 14, no. 16, 15 August 1995, pages 4073-4082, XP002042538 cited in the application  * see the whole document, esp. pp. 4079/80	1-9,27
A	WO 95 07708 A (UNIV CALIFORNIA ;CANJI INC (US)) 23 March 1995 see the whole document	1-29
A	WO 92 05272 A (UNIV CALIFORNIA) 2 April 1992 see the whole document	24,26
A	COLLIN S. ET AL.: "The two nonstructural proteins from wheat dwarf virus involved in viral gene expression and replication are retinoblastoma-binding proteins" VIROLOGY, vol. 219, no. 1, 1 May 1996, pages 324-329, XPO02042539 * see the whole document, esp. p.325, right col. *	1-29
Α	SONI R. ET AL.: "A family of cyclin D homologs from plants differentially controlled by growth regulators and containing the conserved retinoblastoma protein interaction motif" THE PLANT CELL, vol. 7, no. 1, January 1995, pages 85-103, XP002042540 cited in the application * see the whole document, esp. p.97, right col. *	1-29

rational Application No PCT/EP 97/03070

(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/EP 97/03070
tegory °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
,X	XIE Q. ET AL.: "Plant cells contain a novel member of the retinoblastoma family of growth regulatory proteins" THE EMBO JOURNAL, vol. 15, no. 18, 16 September 1996, pages 4900-4908, XP002042541 see the whole document	1-29
X	GRAFI G. ET AL.: "A maize cDNA encoding a member of the retinoblastoma protein family: involvement in endoreduplication" PNAS, U.S.A., vol. 93, no. 17, 20 August 1996, pages 8962-8967, XP002042542 see the whole document	1-29

nternational application No.

PCT/EP 97/03070

Box	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inte	mational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons.
	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
تن ٠٠	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carned out, specifically: Please see Further Information sheet enclosed.
	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6,4(a).
Box II	Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)
This Inter	rnational Searching Authority found multiple inventions in this international application, as follows:
	·
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

### FURTHER INFORMATION CONTINUED FROM PCT/ISAL 10

#### OBSCURITIES:

Claims 28 and 29 are formulated in a very inconcise manner. Consequently, the subject matter claimed was interpreted as follows and searched: Claim 28: "A nucleic acid encoding a protein in which one or more sites are

altered or deleted, making the protein more resistant to the phosphorilation and thus to it's functionality, for example,

linking to E2F or similar "

Claim 29 : Unchanged.

Meaningful search not possible on the basis of all claims: In claim 18 Seq ID 2 was read as Seq ID 1.

Information on patent family members

national Application No PCT/EP 97/03070

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 9506661 A	09-03-95	AU 7642694 A CA 2170605 A CN 1133595 A EP 0716660 A JP 9502183 T ZA 9406595 A	22-03-95 09-03-95 16-10-96 19-06-96 04-03-97 28-02-96
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WO 9205272 A	02-04-92	CA 2091759 A EP 0556231 A JP 6501160 T US 5324654 A	18-03-92 25-08-93 10-02-94 28-06-94